

Drivers and patterns of iron redox cycling from surface to bedrock in a deep tropical forest soil: a new conceptual model

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Abstract Iron (Fe) reduction and oxidation are important biogeochemical processes coupled to decomposition, nutrient cycling, and mineral weathering, but factors controlling their rates and spatial distribution with depth are poorly understood in terrestrial soils. In aquatic ecosystems, Fe reduction often occurs below a zone of oxic sediments. We tested an alternative conceptual model for Fe redox cycling in terrestrial soils using a deep humid tropical forest soil profile. We hypothesized that Fe reduction in

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Department of Environmental Science, Policy, and Management, University of California-Berkeley, Berkeley, CA, USA anaerobic microsites scales with depth variation in labile C and Fe availability, as opposed to bulk oxygen (O_2) . We measured bulk O_2 at multiple depths from 0.1 to 5 m quasi-continuously over 18 months and sampled soils from surface to bedrock (~ 7 m). Median O₂ mixing ratios declined from 19.8 \pm 1.2 % at 0.25 m to 16.1 ± 1.0 % at 1 m, but did not consistently decrease below 1 m, challenging a recent model of regolith development. Reduced Fe (Fe(II)) extractable in 0.5 M hydrochloric acid was greatest in 0-0.1 m soil and declined precipitously with depth, and did not correspond with visible gleying in B horizons. We observed similar depth trends in potential Fe reduction under anaerobic conditions. Depth trends in Fe(II) also closely mirrored short-term soil respiration and bulk soil C. Labile C stimulated Fe reduction at 0-0.1 m depth, whereas addition of short-range-ordered Fe oxides had no effect. Cultivable Fe-reducing bacterial abundance was four orders of magnitude greater in surface soil (0-0.1 m) than below 1 m. Although cultivable Fe oxidizing bacteria were typically also more abundant in surface soil, addition of labile C and nitrate stimulated Fe oxidizers in deep soil by two orders of magnitude under anaerobic conditions. This implies that infiltration of nitrate (and possibly C) from shallow soil water could potentially promote biotic Fe oxidation, a critical step in bedrock weathering, 7 m below. Together, these data suggest that C, Fe, and nutrient availability increase microbial Fe reduction and oxidation in surface (vs deeper) soil microsites despite high bulk O₂, in contrast to the depth segregation of electron accepting processes often observed in aquatic ecosystems. Furthermore, the greatest capacity for Fe redox cycling can occur in A horizons that do not display gleying or mottling.

Keywords Critical zone · Iron oxidation · Iron reduction · Mineral weathering · Oxygen · Redox · TEAP

Introduction

Iron (Fe) oxidation and reduction driven by microbial and/or abiotic processes are coupled to the biogeochemical cycling of carbon (C), phosphorus (P), nitrogen (N), and cations over ecological timescales, and contribute to mineral weathering and soil evolution over pedogenic timescales. Dissimilatory Fe reduction coupled to C oxidation is an important anaerobic microbial respiratory process, and dark Fe oxidation coupled to oxygen (O_2) or nitrate (NO_3^-) reduction can also support microbial growth (Weber et al. 2006; Melton et al. 2014). The ecosystem-scale importance of Fe redox cycling and its relationships to other elemental cycles have received greatest attention in aquatic sediments and wetland soils (Ponnamperuma 1972; Lovley 1995; Thamdrup 2000; Weber et al. 2006; Cheng et al. 2010). Yet, Fe redox cycling can also influence organic matter decomposition, nutrient dynamics, and mineral weathering in relatively well-drained surface soils of terrestrial ecosystems (Chacón et al. 2006; Thompson et al. 2006; Fimmen et al. 2008; Dubinsky et al. 2010; Hall and Silver 2013; Yang and Liptzin 2015). These dynamics are especially relevant in humid tropical soils, which are often rich in short-range-ordered Fe oxides and organic C. In these ecosystems, rates of soil Fe redox cycling and pools of reduced Fe (Fe(II)) often equal or exceed wetland sediments (Dubinsky et al. 2010; Thompson et al. 2011; Hall and Silver 2015). Yet, understanding the spatial distribution and controls on Fe reduction and oxidation in terrestrial soils remains an important knowledge gap hampering the incorporation of Fe redox cycling into quantitative and conceptual models of short term (i.e., minutesmonths) ecosystem dynamics, and long-term (i.e., centennial-millennial) weathering and pedogenic processes.

In aquatic sediments and groundwater, a dominant conceptual model proposes that respiratory terminal electron accepting processes exhibit an approximate segregation with depth according to their thermodynamic favorability. That is, given sufficient supply of C or other reductants, O₂ is rapidly reduced near the sediment surface, followed by the reduction of nitrate, manganese oxides, and Fe oxides in progressively deeper zones (Froelich et al. 1979; Chapelle et al. 1995; Roden and Wetzel 1996; Emerson and Hedges 2003). Thus, in undisturbed sediments we would typically expect respiratory Fe reduction to commence at depths where most O_2 has been depleted (Fig. 1a) although sediment redox gradients can also be disrupted by bioturbation (Norkko et al. 2011). In terrestrial environments, spatial and temporal heterogeneity in O_2 availability is a common feature of soils undergoing fluctuations in moisture, C inputs, and biological activity. The importance of microsite-scale (mm-cm) redox gradients for stimulating denitrification in well-drained surface soils is widely



Fig. 1 Idealized depth profiles of bulk O_2 (*solid line*) and Fe(II) (*dashed line*) with depth. *Panel* **a** represents a standard conceptual model for aquatic sediments, where Fe reduction occurs at depths below which O_2 and other oxidants (not shown for clarity) have been reduced. *Panel* **b** shows an alternative model for terrestrial soils where bulk O_2 decreases slightly with depth, and total Fe reduction peaks in anaerobic microsites near the soil surface as a consequence of greater availability of C and short-range-ordered Fe. Oxygen profiles were modeled after Cerling (1991), assuming a stoichiometric relationship between CO_2 and O_2 and an exponential decline in CO_2 production with depth; diffusivity differs for *panels* **a** and **b**. Iron(II) trends are hypothetical but consistent with previous work at this site; labile Fe(II) may increase near bedrock due to supply from primary minerals (Buss et al. 2005)

acknowledged, and has been contrasted with the depth stratification of redox reactions in aquatic sediments (Seitzinger et al. 2006). Iron reduction also appears to be relatively commonplace in many well-drained surface soils (Silver et al. 1999; DeAngelis et al. 2010; Liptzin et al. 2011; Yang and Liptzin 2015), and even gross methane production can sometimes be measured in these systems (von Fischer and Hedin 2007). Despite these observations, patterns in the depth distribution of Fe redox cycling vis a vis availability of O_2 and other potential drivers (C, Fe, and N) have received much less attention in terrestrial soils.

Spatial interactions between controls on physical O₂ supply and biological O₂ demand may be crucial for understanding trends in Fe redox cycling with depth in terrestrial soils. Macropore carbon dioxide (CO_2) concentrations typically increase with soil depth (Cerling 1991), corresponding to a stoichiometric decrease in O2 that could stimulate Fe reduction with depth. Indeed, several studies have documented Fe reduction in subsoils (0.5-1.5 m) using morphological observations and geochemical analyses (Veneman et al. 1976; Fimmen et al. 2008; Schulz et al. 2016). The combination of periodically perched water and/or root C inputs to clay-rich subsurface horizons appeared to promote Fe reduction and oxidation in these studies (ibid.), generating prominent visual features of gleying and mottling indicative of Fe redox cycling. However, significant rates of Fe reduction can also occur in surface (A) soil horizons from a broad range of ecosystems (Chacón et al. 2006; Thompson et al. 2006; Dubinsky et al. 2010; Buettner et al. 2014; Yang and Liptzin 2015). At the surface, development of aggregates with tortuous diffusion paths allows anaerobic processes to occur in close spatial proximity to macropores with near-atmospheric O₂ concentrations (Sexstone et al. 1985). As a consequence, measurements of O₂ in soil macropores (defined here as "bulk O₂") do not necessarily reflect the prevalence of anaerobic microsites at small (mm-cm) spatial scales, despite their utility when comparing among sites over larger (m-km) spatial scales (Silver et al. 2013; Hall and Silver 2015; Liptzin and Silver 2015). Because both the availability of O_2 in soil macropores as well as total biological O_2 demand generally decrease with depth (Cerling 1991), the overall relationship between soil depth and Fe reduction remains unclear.

The availability of short-range-ordered Fe oxides, organic C, and co-limiting nutrients could also have a crucial impact on the depth distribution of Fe reduction. Humid tropical soils are often rich in Fe oxides, especially goethite and hematite, as a consequence of extensive weathering and desilication (Sanchez 1976; White et al. 1998). Yet, a relatively small fraction of total Fe may be readily accessible to Fe-reducing microbes. Iron reduction rates often scale with the surface area and solubility of Fe oxide phases (Roden and Zachara 1996; Bonneville et al. 2009). The shortrange-ordered Fe phases that dominate reducible Fe pools (Hyacinthe et al. 2006) may decline with depth (Thompson et al. 2011; Hall and Silver 2015), potentially limiting Fe reduction. Organic C availability may also limit Fe reduction with depth. Humid tropical forests are characterized by high C availability that fuels heterotrophic activity in surface soils (Raich and Schlesinger 1992), and rates of Fe reduction appear tightly coupled with the availability of dissolved organic C (Chacón et al. 2006; Fuss et al. 2010). Other nutrients such as nitrogen (N) could also limit Fe reduction/oxidation, especially in deeper soil horizons with low organic matter content. Even in comparatively N-rich tropical forests, N additions can enhance particulate organic matter decomposition (Cleveland and Townsend 2006; Cusack et al. 2011), and nutrient limitation may be exacerbated in comparatively resource-poor subsoils (Stone et al. 2014).

In deep soils, Fe(II) oxidation is also a crucial step in bedrock weathering, where minerals such as hornblende and biotite provide a source of Fe(II) that can be oxidized via biotic or abiotic mechanisms coupled to O_2 or NO_3^- (Buss et al. 2005; Fletcher et al. 2006; Liermann et al. 2015). It has been hypothesized that O₂ availability limits mineral weathering at the interface between bedrock and saprolite, and thus may play a key role in controlling landscape evolution (Fletcher et al. 2006; Brantley and White 2009; Bazilevskaya et al. 2013; Behrens et al. 2015). Oxygen concentrations often decrease with depth, reflecting a balance between diffusive supply and the biological and geochemical processes that consume O_2 . Fletcher et al. (2006) proposed that depth-dependent decreases in O_2 served as a negative feedback on bedrock weathering, given that increasing regolith thickness would presumably result in decreased O₂ supply at the weathering front. Alternative oxidants such as NO₃⁻ could also potentially contribute to Fe(II) oxidation

(Böhlke et al. 2002; Liermann et al. 2015), especially under O_2 -limited conditions.

Few studies have examined trends in O_2 and biogeochemical processes across deep soil profiles. In Amazonian forests and pastures, soil CO₂ concentrations increased monotonically with soil depth, implying a corresponding stoichiometric decline in O_2 from ~19 % above 1 m to ~12 % at 8 m (Nepstad et al. 1994). Similarly, in a highly weathered Puerto Rican forest soil, O_2 declined from ~18 % above 2 m to ~13 % at 7 m (Liermann et al. 2015). It is unclear whether changes in O_2 availability of this magnitude might impact biotic Fe(II) oxidation at depth, and whether Fe-oxidizing microbial abundance might respond to availability of O_2 , NO_3^- , or organic matter.

We tested the hypothesis that microbial capacity for Fe reduction and oxidation across a deep tropical forest soil profile correlates with depth variation in the availability of C and short-range-ordered Fe as opposed to bulk soil O₂. In accordance with this hypothesis, we predicted greater rates of Fe reduction and oxidation potential in surface as opposed to deeper soils, despite a predicted decline in bulk soil O₂ with depth (Fig. 1b). This conceptual framework contrasts with patterns often observed in flooded wetland soils and sediments (Fig. 1a). We predicted that our terrestrial soils would deviate from this spatial segregation of aerobic and anaerobic processes with depth, because of the importance of electron donor supply (i.e., organic C) in generating anaerobic microsites where Fe reduction can occur within a porous soil matrix, as well as an increased abundance of shortrange-ordered Fe in surface soils. We also assessed the degree to which addition of C, Fe, and NO₃⁻ affected Fe reduction and oxidation capacity across the depth gradient. We predicted that NO_3^- addition would increase Fe oxidation capacity in deep soils, where bedrock supplies ample Fe(II) but oxidant (O₂, NO₃⁻) availability may limit rates of Fe(II) oxidation.

Methods

Site description

Samples were collected from the Guaba Ridge (18°17′02″N, 65°47′20″W) in the Río Icacos Watershed of the Luquillo Experimental Forest, Puerto Rico.

This humid montane tropical forest ecosystem has mean annual temperature and precipitation of 22 °C and 4200 mm, respectively (White et al. 1998). Despite high precipitation, surface soils (0–10 cm) remain well drained due to high porosity (\sim 75 %; White et al. 1998) and bioturbation. Parent material is quartz diorite from the Rio Blanco stock, dominated by plagioclase feldspar and quartz (White et al. 1998). Soils in the watershed include Oxisols, Ultisols, and Inceptisols, and vary according to topographic position (Soil Survey Staff 2002; Johnson et al. 2015). The Guaba Ridge separates two first-order streams that discharge to the Río Icacos at approximately 650 m elevation. The soil sampled here was recently characterized as a Plinthic Haplohumult (Yi-Balan et al. 2014), similar to the Humic Hapludox described by the Soil Survey Staff (2002). The B horizons transition to saprolite at a depth of approximately 1 m (White et al. 1998; Yi-Balan et al. 2014). Gleying indicative of Fe reduction was especially prominent between 0.2 and 0.4 m. Fine root biomass was greatest from 0 to 10 cm and declined precipitously with depth, and was absent below 80 cm (Johnson et al. 2015; Hall and Silver 2015). Organic C declined with depth from 2 to 3 % C by mass from 0 to 0.1 m (S. J. Hall, unpublished data), to 1.6, 1.5, 1.2, and 1.1 % C at depths of 0.15, 0.3, 0.45, and 0.6 m, respectively. Below 0.6 m, C was typically <0.2 % (Buss et al. 2005). Clay-sized particles were most abundant (42 %) at 0.3 m, and measured between 16 and 30 % in other samples to 5 m depth (Buss et al. 2005). Total Fe oxide content (as Fe₂O₃) increased from ~ 4 % at the surface to >7 % at depth (White et al. 1998). Site vegetation was evergreen tropical montane forest locally described as the "palo colorado" forest, after the dominant species Cyrilla racemiflora L. (Weaver and Murphy 1990).

Soil sampling

Soils were sampled on two separate occasions using a 7.6 cm diameter stainless steel bucket auger and extensions. Samples at a given depth interval represent composites from three separate augured holes collected within a radius of 15 m. The 2010 samples were collected from depth increments of 0–0.15, 1.5–1.8, and 6.9–7.2 m. The 2012 samples were collected from 0 to 0.1, 0.1 to 0.2, 0.2 to 0.5, 0.5 to 1, 1 to 2, 2 to 3, 3 to 4, and 4 to 5 m. Samples were stored at field moisture

in sealed polyethylene bags at ambient temperature (22-25 °C).

Oxygen measurements

We installed O₂ sensors (Apogee SO-110, Logan UT) at depths of 0.1, 0.25, 0.5, 1, 2, 3, 4, and 5 m in June 2010 and monitored them until February 2012. Sensors were calibrated at 100 % relative humidity prior to installation and upon retrieval, and corrected for linear drift over time. Each sensor was installed in a separate hole augured to the depth of installation. Holes were separated laterally by >1 m. Sensors were deployed inside 10 cm lengths of 5.1 cm diameter polyvinylchloride pipe sealed with a cap on the top and bottom and perforated on the sides with 0.5 cm diameter holes to allow gas exchange with the adjacent soil atmosphere. After lowering a sensor to the bottom of the augured hole, soil was refilled and tamped above the sensor to approximate field bulk density using a stainless steel rod. The initial week of data was discarded, after which point O₂ concentrations (atmospheric mixing ratios) established pseudo steady-state values at deeper depths. Data were recorded at hourly intervals on a datalogger (CR1000, Campbell Scientific, Logan UT) during most of the 21-month period. Continuous measurements were not possible due to remote nature of the field site and associated battery failure.

Chemical analyses and laboratory experiments

We measured Fe and trace gas production in the laboratory at U. C. Berkeley shortly after soils were sampled, and during the course of two laboratory experiments. Iron(II) and (III) were measured in 0.5 M HCl extractions using a 1:10 mass ratio of soil to solution, denoted as Fe(II)_{HCl} and Fe(III)_{HCl}. Soils were extracted for 2 h on a rotary shaker, centrifuged at 3200 rcf, and the supernatant solution filtered to 0.2 µm. Solutions were analyzed using a modified ferrozine method (Viollier et al. 2000). Here, we used Fe(III)_{HCl} as an index of short-range-ordered Fe oxides. Our previous work at nearby sites showed a strong correlation between Fe(III)_{HCI} and Fe extracted via reductive dissolution with citrate-ascorbate solution, although Fe(III)_{HCl} was always of smaller magnitude (Hall and Silver 2015). Citrate-ascorbate extractable Fe is thought to be closely correlated with microbially-reducible Fe (Hyacinthe et al. 2006). We measured production of carbon dioxide (CO_2) using gas chromatography (Shimadzu 14A, Columbia MD) as described previously (Hall et al. 2013).

We tested relationships between bulk soil O₂, trace gas production, and Fe_{HCl} across soils from 0 to 5 m depth (0–0.1, 0.1–0.2, 0.2–0.5, 0.5–1, 1–2, 2–3, 3–4, and 4–5 m). Samples (~15 g dry mass equivalent) were incubated in glass jars for 24 h in darkness under an ambient atmosphere (20.9 % O₂) within 7 days of sample collection, with three replicates per depth. We report Fe(II)_{HCl} and Fe(III)_{HCl} extracted immediately prior to the trace gas measurements. Next, we incubated a subset of these soils (0.1–0.2, 0.5–1, 2–3, 3–4, 4–5 m depths) under hypoxic (N₂ headspace) and aerobic conditions (~20.9 % O₂) over 10-days to assess potential rates of Fe reduction (n = 3 per depth and headspace).

Together, these measurements identified surface soil horizons as a dominant zone of actual and potential Fe reduction. We then tested the importance of labile C and short-range-ordered Fe availability as controls on Fe reduction in 0-0.1 m soil using a full factorial experiment (n = 3 per treatment) conducted under hypoxic conditions (N2 headspace) to simulate the presence of reducing microsites under field conditions. Short-range-ordered Fe as hydrous ferric oxide was prepared as described previously (Yang et al. 2012) and gently homogenized with soil subsamples (~ 15 g dry mass equivalent) at concentrations of 0, 0.1, 0.5, and 1 mg Fe g soil⁻¹. Labile C was added as glucose dissolved in deionized water at concentrations of 0, 50, 100, and 200 μ g C g soil⁻¹. Glucose was used given that it can be fermented to multiple compounds that support Fe reduction (Lovley 1995).

Finally, we used separate samples spanning the soil surface to bedrock to further test environmental controls on Fe reduction, as well as the abundance of Fe reducing and oxidizing bacteria using most probable number (MPN) analyses as described by Dubinsky et al. (2010). Soils from 0 to 0.15, 1.5 to 1.8, and 6.9 to 7.2 m depths were combined in a 1:2 ratio with deionized water and incubated under anaerobic conditions for 8 days. This experiment was designed to assess controls on Fe cycling under anaerobic conditions, to test impacts of NO_3^- availability on Fe oxidation, and to compare with an aerobic pretreatment control. Soil solutions were amended with

either sodium nitrate (NO₃⁻; 1 mM final concentration), Fe as ferrous chloride (2 mM), Fe+NO₃⁻, sodium acetate (0.5 mM), and acetate+ NO_3^- , or deionized water, and incubated in an anaerobic chamber (90 % N₂, 8 % CO₂, and 2 % H₂ headspace). Acetate was used in this experiment given the precedence of studies that successfully cultivated Fe oxidizers and reducers (Lovley 1995; Straub et al. 1996). To enumerate anaerobic Fe reducing and oxidizing bacteria, soils were extracted in buffer containing 0.1 % sodium pyrophosphate and 0.03 % Tween 80 in basal microbiological medium (BMM). BMM consisted of (per L) 5.0 g 2-(N-morpholino)ethanesulfonic acid (MES) buffer and 10 ml mineral solution, with 0.80 g NaCl, 1.0 g NH_4Cl , 0.1 g KCl, 0.1 g KH₂PO₄, 0.2 g MgCl₂·6H₂O, and 0.04 g CaCl₂·2H₂O (per L). After autoclaving, the media pH was adjusted to 5.5 and amended with 1 ml SL12 trace elements solution, 2.5 ml trace metal solution (Widdel and Bak 1992), and 1 ml vitamin solution (Pfennig and Trüper 1992) per L. Media was dispensed into 96-well microplates, and soil subsamples added in ten-fold dilutions from 10^{-2} to 10^{-13} with four biological replicates and three technical replicates per depth per amendment. Plates were incubated in the dark for 30 days with negative controls including soil extract buffer only (no soil). Positive growth of Fe(III) reducers was visualized by adding ferrozine solution, which turns purple in the presence of Fe(II). Formation of reddish-brown precipitates was used to verify positive results for Fe(II) oxidizers. Cell counts per gram of soil were calculated using the MPN Calculator version 4.04 (Klee 1996).

For all experiments, statistical differences among treatments and/or depths were assessed using ANOVA and post hoc Tukey comparisons using R v. 3.2.0. To account for temporal autocorrelation in the O_2 data, we used a generalized linear model with an autoregressive error term implemented using the glm function.

Results

often small: below 0.5 m, median O_2 concentrations differed by <0.3 % and did not consistently decrease with depth (Fig. 2a). Oxygen was most dynamic at 0.25 and 0.5 m depths, where O_2 varied by as much as 10 % over time (Fig. 2a). In contrast, depths below 0.5 m showed much less variability (<1.5 % O_2) relative to median values.

Concentrations of Fe(II) HCl and Fe(III)HCl measured on soils sampled in May 2012 showed different patterns from bulk soil O₂ (Fig. 2). Iron(II)_{HCl} was greatest in 0-0.1 m soil and declined precipitously with depth, and was negligible below 1 m (Fig. 2b). Concentrations of Fe(III)_{HCl} showed very similar depth trends as Fe(II)_{HCl} (Fig. 2c). Patterns of shortterm CO₂ production closely mirrored both Fe(II)_{HCl} and Fe(III)_{HCl} (Fig. 2d), and rates declined >tenfold between the 0-0.1 and 0.5-1 m depths. Ten-day anaerobic incubations of a subset of these soils confirmed that potential Fe reduction was greatest in the most shallow soil tested (0.1-0.2 m), declined by an order of magnitude between 0.5 and 3 m, and was undetectable (not different from zero, p > 0.05) from 3 to 5 m (Fig. 3). Samples incubated under an *aerobic* atmosphere displayed no significant net change in Fe(II) concentrations over this time period (data not shown).

Factorial incubation experiments with 0-0.1 m soil indicated that Fe reduction generally increased with increased rates of labile C addition (glucose) under anaerobic conditions. Production of Fe(II)_{HCl} was significantly greater (p < 0.05) in samples that received the highest glucose concentrations (100 and 200 μ g C g soil⁻¹) and no or minimal Fe(III) addition (0 and 0.1 mg Fe g soil⁻¹). Iron(III) addition had no significant effect on Fe(II)_{HCl} production in the treatments with no or minimal glucose addition (0 and 50 μ g C g soil⁻¹). However, the treatments with the highest Fe concentrations decreased Fe(II)_{HCl} production relative to the controls. This effect depended on the amount of added glucose (Fig. 4; treatment interaction p < 0.0001). Addition of 1 mg Fe g soil⁻¹ decreased Fe(II)_{HCl} production in the presence of 100 μ g glucose C g soil⁻¹, whereas 0.5 mg Fe g soil⁻¹ decreased $Fe(II)_{HCl}$ production with 200 μ g glucose C g soil⁻¹.

In our final experiment, we incubated soil slurries from surface, intermediate, and deep samples (0-0.15, 1.5-1.8, and 6.9-7.2 m) to test the factors controlling the abundance of Fe reducing and oxidizing organisms



Fig. 2 Soil O_2 concentrations a measured at hourly intervals along the Guaba ridge depth profile from June 2010 to February 2012. *Boxes* represent medians and the first and third quartiles. *Whiskers* represent the furthest value less than 1.5 times the box length measured from the box edge; more extreme points are



Fig. 3 Net Fe(II) production by depth $(\pm SE)$ over a 10-day incubation of intact (non-slurried) soils under anaerobic conditions (n = 3 per depth). Note that *bar* widths are not proportional to soil depths (as in Fig. 2) because a subset of depths was measured

under anaerobic conditions, simulating anaerobic microsites in the field. Iron reduction rates were two orders of magnitude greater in surface than deeper samples (p < 0.001), although lower but significant rates of Fe reduction were also detectable in some of the intermediate and deep samples under these experimental conditions (Fig. 5a, note the log scale). Trends in Fe reduction rates with depth were corroborated by

denoted as *circles*. Mean (±SE) concentrations of Fe(II)_{HCI} **b** Fe(III)_{HCI} **c** and soil respiration **d** were measured shortly after sampling (n = 3 per depth). Means with *different letters* differed significantly (p < 0.05, Tukey comparison)



Fig. 4 Iron reduction during three-day anaerobic incubations of intact (non-slurried) soils from 0 to 0.1 m depth, incubated with varying concentrations of labile C (glucose) and short-range-ordered Fe(III). Treatments with *different letters* differed significantly (all possible pairwise comparisons were evaluated), and *whiskers* represent standard errors (n = 3 per treatment)

MPN analyses of cultivable Fe reducers, which were four orders of magnitude greater in surface than deeper samples (Fig. 5b). Experimental amendments (acetate, Fe, and NO_3^-) did not significantly affect Fe reduction rates in surface (0–0.15 m) samples. However, Fe + NO_3^- addition increased the abundance of



Fig. 5 Rates of Fe reduction (a) and most-probable-number (MPN) analyses of Fe reducers (b) and oxidizers (c) measured before and after anaerobic incubations of soil slurries. Samples were amended with NO_3^- , Fe as ferrous chloride, NO_3^- + Fe,

cultivable Fe reducers three-fold relative to the controls in these samples (p < 0.05; Fig. 5b). Nitrate addition alone doubled mean cultivable Fe reducer abundance relative to the control in surface soil, but this difference was not statistically significant. In the intermediate depth samples (1.5-1.8 m), addition of Fe, Fe + NO_3^- , acetate, and acetate + NO_3^- all significantly stimulated Fe reduction rates relative to the control, but had no significant impact on Fe reducer MPN (Fig. 5a, b), where abundances were low across all treatments. Iron $+ NO_3^-$ addition stimulated Fe reduction rates in the intermediate depth samples to the greatest extent (p < 0.05). In the deep soil samples (6.9–7.2 m), Fe and Fe + NO_3^- stimulated Fe reduction relative to the controls, whereas the other treatments had no significant effects.

Iron(II) oxidizers were most abundant in surface soils at the beginning of the experiment, and Fe oxidizer MPN was not significantly affected by experimental amendments in either the surface or intermediate samples (Fig. 5c). In the deep samples, addition of acetate $+ NO_3^-$ (but not acetate or $NO_3^$ alone) significantly stimulated Fe oxidation MPN by two orders of magnitude relative to the other treatments. Notably, Fe oxidizer MPN values were not inhibited by anaerobiosis per se at any depth, as they

given depth increment differed significantly (n = 4 per

inhibited by anaerobiosis per se at any depth, as they did not decrease relative to initial values under any treatment despite incubation under anaerobic conditions.

Discussion

treatment)

Our data support the hypothesis that labile C and Fe availability (Fig. 1b), as opposed to variation in bulk O₂ with depth (Figs. 1a, 2), controlled Fe reduction across this deep humid tropical forest soil profile. Actual and potential Fe reduction and cultivable Fe reducer abundance were greatest in surface soils where bulk O₂ concentrations were also highest (Figs. 2, 3). Iron reduction declined by two orders of magnitude below 1 m despite decreased bulk soil O₂, and cultivable Fe reducer abundance declined by four orders of magnitude. Soil respiration (Fig. 2) and bulk soil C (Buss et al. 2005) showed a depth pattern similar to Fe reduction, and addition of labile C (but not Fe) significantly enhanced Fe reduction in surface soil (Fig. 4). Apparent decreases in $Fe(II)_{HCI}$ production at the highest concentrations of added Fe and glucose (Fig. 4) may represent Fe(II)-catalyzed transformation

of short-range-ordered Fe into more crystalline phases that occluded the newly-produced Fe(II) (Jeon et al. 2003). Although these data represent a single site, they support our proposed conceptual framework as well as the need to more broadly reconsider the controls and impacts of Fe redox cycling with depth, as discussed below.

Contrasting depth distribution of Fe redox cycling in wetlands and uplands

A dominant conceptual model in aquatic sediments posits that terminal electron accepting processes are segregated with depth according to their thermodynamic favorability (Emerson and Hedges 2003; Fig. 1a). However, our data suggest that this model does not necessarily explain depth variation in Fe reduction and oxidation potential at our site, and perhaps also in other well-drained terrestrial soils where high Fe reduction capacity has been documented (Yang and Liptzin 2015). Bulk O₂, Fe(II) concentrations, and potential Fe reduction were all greatest at the surface, concomitant with greatest labile C availability. These observations, combined with our experimental data, suggest that patterns in electron donor (i.e., organic C) availability provide a proximate control on the depth distribution of Fe reduction in this soil.

Although bulk O2 concentrations declined with depth below 0.25 m, the abundance of anaerobic microsites also declined along with C availability, as reflected by lower concentrations of Fe(II), lower potential Fe reduction rates, and lower abundance of Fe-reducing and oxidizing microbes. Subtly increased O₂ concentrations at 0.25 m relative to 0.1 m likely reflected lateral spatial heterogeneity in soil O₂ (Liptzin et al. 2011; Hall et al. 2013) as opposed to a consistent trend with depth. Contrasting patterns between the depth distribution of terminal electron accepting processes in aquatic sediments versus this terrestrial forest point to the importance of micrositescale anaerobic processes within the largely aerobic soil matrix (Sexstone et al. 1985; Hall and Silver 2015; Keiluweit et al. 2016). Localized inputs of labile C to fuel O2 consumption may be a critical regulator of the abundance of anaerobic microsites (Chacón et al. 2006), evidenced by recent reports of Fe reduction associated with the rhizosphere (Fimmen et al. 2008; Schulz et al. 2016). In many ecosystems, including this humid tropical forest, root biomass C inputs are greatest near the soil surface (Jobbagy and Jackson 2000; Hall and Silver 2015). Thus, we predict that Fe redox cycling and the numerous processes linked to these dynamics—e.g., sorption and desorption of P and organic matter (Peretyazhko and Sposito 2005; Chacón et al. 2006; Thompson et al. 2006; Buettner et al. 2014; Hall et al. 2016), microbial respiration (Dubinsky et al. 2010), and production of reactive oxygen species (Hall and Silver 2013)—may also be most significant near the surface of many other terrestrial soils, despite the fact that moisture and visual indicators of Fe redox cycling often increase with depth.

Cryptic Fe redox cycling in surface soils

Our finding of greater Fe reduction capacity in surface versus subsurface soils shows that trends in soil coloration and moisture are not necessarily reliable indicators of potential rates of Fe redox cycling. The greatest Fe(II) concentrations, potential rates of Fe reduction, and abundance of cultivable microbial Fe reducers and oxidizers occurred in surface A horizon (0-0.1 m) soil, where porosity was high (~75 %) and moisture rarely approached saturation (White et al. 1998). In this soil, gleying was visible throughout the B horizons, but not in the A horizon (Yi-Balan et al. 2014). Surface horizons were rarely saturated, and water content was typically greatest near the soilsaprolite interface (White et al. 1998). Investigations of Fe reduction in terrestrial soils have often focused on Fe redox dynamics in relatively deeper (>0.5 m) B horizons associated with periodic moisture saturation and rhizosphere gleying (Veneman et al. 1976; Fimmen et al. 2008; Schulz et al. 2016). Mottling and gleying provide important visual evidence of Fe reduction, translocation, and oxidation (Veneman et al. 1976; Fimmen et al. 2008; Schulz et al. 2016). Although these features are *sufficient* to indicate the occurrence of Fe redox cycling, they are not obligate indicators, as demonstrated by our data. Several studies have similarly demonstrated high rates of Fe reduction at the soil surface (Chacón et al. 2006; Dubinsky et al. 2010; Yang and Liptzin 2015) but did not assess trends with depth. Visual evidence of Fe redox cycling in 0-0.1 m soil may have been obscured by high organic matter content at the surface, which imparts a dark color. Surface soil horizons typically

contain the highest stocks of root biomass and organic C across a broad range of terrestrial ecosystems (Jobbagy and Jackson 2000). As a consequence of abundant C inputs that generate anaerobic microsites yet obscure the visual effects of Fe reduction, we suggest that cryptic Fe reduction in terrestrial surface soils may be a more commonplace phenomenon than is implied by visible gleying and mottling.

In surface soils of humid tropical forests and other terrestrial ecosystems, hotspots of Fe reduction in surface soil microsites are likely generated due to the confluence of several critical factors. High clay content decreases gas-phase diffusivity and O₂ supply, high temperature decreases O2 solubility while increasing biological O₂ consumption, a high density of live and dead roots provides abundant C inputs, and large pools of Fe are maintained in short-rangeordered minerals (Silver et al. 1999; Buss et al. 2005; Thompson et al. 2011; Johnson et al. 2015; Hall and Silver 2015). The maintenance of short-range-ordered Fe appears critical in that these phases exhibit greater rates of reduction than crystalline Fe (Roden and Wetzel 2002). Interactions between Fe and organic matter likely retard the formation of crystalline minerals (Schwertmann et al. 1988), despite the fact that redox cycling can potentially lead to formation of Fe with greater crystalline structure (Thompson et al. 2006). At depth, lower organic matter concentrations may facilitate the formation of more crystalline Fe minerals during redox cycling (Jeon et al. 2003; Thompson et al. 2006), consistent with previous Fe isotope measurements at this site (Buss et al. 2010). This coincides with our finding that Fe addition stimulated Fe reduction at depth (Fig. 5), despite the presence of a large total Fe oxide pool (White et al. 1998).

Implications of O₂ depth distribution for bedrock weathering

Bulk soil O_2 concentrations did not consistently decrease with depth below 1 m, the approximate depth of the soil/saprolite transition, but rather fluctuated around similar median values (~16 %; Fig. 2). Previous work hypothesized that regolith depth controls weathering rates by constraining O_2 supply, given that O_2 availability may limit oxidation of Fe in primary bedrock minerals (Fletcher et al. 2006; Brantley and White 2009; Behrens et al. 2015). A key assumption of this hypothesis is that O₂ concentrations decrease monotonically with regolith depth, facilitating a negative feedback between regolith development and weathering rates. Our data show that bulk soil O2 does not necessarily decrease consistently or significantly within the saprolite profile. Rather, the asymptotic trend in bulk soil O₂ observed here at depths below 1 m is consistent with analytical models of soil CO₂ production and diffusion (Cerling 1991; Fig. 1b) validated by measurements in shallower soil profiles (<1.5 m depth) from other ecosystems (e.g. Solomon and Cerling 1987; Bowling et al. 2015). Bulk O_2 concentrations observed here at 5 m depth (\sim 16 %), as well as other tropical forests $(\sim 12-17 \%$; Nepstad et al. 1994; Liermann et al. 2015), were relatively high. If these data and models are broadly representative, regolith depth per se may not necessarily influence gas-phase O₂ supply to the bedrock/saprolite interface. Rather, the presence and depth of perched water tables at the bedrock/saprolite interface (White et al. 1998) may be more important in controlling diffusive O₂ supply for primary mineral weathering.

Potential importance of anaerobic Fe oxidation at depth

Although bulk soil O₂ concentrations were relatively high in deep soils, anaerobic microsites are likely to occur (Silver et al. 1999), particularly in the presence of perched water tables at the bedrock interface (White et al. 1998; Schulze and White 1999). Iron(II) oxidation by NO₃⁻ can be a significant process in shallow groundwater (Böhlke et al. 2002). As a consequence, availability of both O2 and NO3⁻ could potentially influence rates of Fe(II) oxidation at the bedrock/saprolite weathering front. In our study, Fe(II) oxidation at intermediate depths was likely limited by Fe(II) supply from primary minerals (Buss et al. 2005) and the abundance of cultivable Fe(II) oxidizers was unaffected by our experimental treatments. In deep soils, cultivable Fe(II) oxidizer abundance was initially similar to intermediate depths, but responded strongly to additions of NO_3^- and acetate under anaerobic conditions-increasing by two orders of magnitude relative to initial aerobic conditions. This finding suggests significant capacity for anaerobic, microbially-mediated Fe(II) oxidation in deep soils.

We note that our results are likely conservative in that the MPN enumeration method used here yielded *cultivable* anaerobic Fe reducers and oxidizers, and likely underestimates their total populations (Dubinsky et al. 2010). Previous work in nearby soils found that cultivable Fe reducers represented 0.7–5.7 % of total bacterial abundance, but that the relative abundances of the canonical Fe reducers *Shewanella* and *Geobacter* assessed were low when assessed using quantitative PCR (Dubinsky et al. 2010; DeAngelis et al. 2010). The composition of Fe reducing and oxidizing microbial communities in humid tropical soils remains poorly understood.

The finding that acetate stimulates anaerobic Fe(II) oxidizers suggests that heterotrophic or mixotrophic Fe oxidizers contribute to Fe(II) oxidation and related bedrock weathering at depth. Previous studies similarly found that C addition enhanced rates of Fe(II) oxidation (Straub et al. 1996; Kappler et al. 2005). The stimulatory effect of acetate on Fe(II) oxidizer abundance may reflect the importance of mixotrophy in preventing the deleterious effects of cell encrustation by the newly-formed Fe(III) oxides (Kappler et al. 2005). However, the finding that acetate stimulated Fe(II) oxidizer abundance presents an interesting conundrum: previous measurements suggested that deep dissolved organic C concentrations may be extremely low at this site (Schulz and White 1999), supporting the hypothesis that microbial communities are dominated by autotrophs reliant on bedrock Fe(II) supply and are decoupled from surface C inputs (Liermann et al. 2015). Yet, the strong response of cultivable Fe oxidizers to C addition in deep soils also suggests that heterotrophic microbial communities are poised to respond to C inputs, either from co-occurring autotrophs or possibly from surface soils.

The observation of high NO₃⁻ concentrations (~20 μ M) in deep soils from this site and another nearby site (Schulz and White 1999; Liermann et al. 2015) is indicative of hydrologic NO₃⁻ supply from surface soils 7 m above, as the parent material does not contain significant N (White et al. 1998). Surface soil biological processes do not appear strongly N limited in this ecosystem (Cusack et al. 2011), thus infiltration of surface soil NO₃⁻ to deep soils appears plausible. The potential for surface soil dissolved organic matter to reach the weathering front at 7 m depth without being sorbed or mineralized in transit may be more tenuous. Couplings between surface-

derived nutrients and bedrock weathering remain an important but poorly explored topic in the context of landscape evolution, and biogeochemical connections between surface and deep subsurface soils merit further exploration.

Conclusions

Trends in potential Fe reduction and oxidation varied systematically with depth in this terrestrial humid tropical forest soil but showed distinctly different trends compared to the standard conceptual model for saturated sediments. Although mean bulk O2 declined overall with depth, it was most variable and sporadically reached the lowest values at shallow depths (0.25-0.5 m). Biotic Fe reduction and oxidation capacity were greatest at the surface and declined precipitously with depth. At intermediate and deep depths, Fe reduction and oxidation appeared strongly limited by C, NO₃⁻, and/or labile Fe, despite high total Fe. However, biotic Fe oxidation potential increased at the saprolite/bedrock interface in response to acetate and NO_3^- addition, likely as a consequence of increased Fe(II) supply from primary Fe(II)-rich minerals, which had been depleted from shallower saprolite (intermediate depths). Shallow surface soils may play an underappreciated role as hotspots of coupled Fe reduction and oxidation, even when visible gleying is not apparent. Furthermore, our data suggest that the total depth of soil profiles may have less influence on bulk O₂ supply to bedrock than previously proposed, given the observed asymptotic trend in O_2 with depth. In addition to O_2 availability, we showed that the supply of NO₃⁻ from surface soils could play an important role in bedrock weathering by stimulating Fe(II) oxidizing microbial communities. Although labile C amendments stimulated the growth of Fe(II) oxidizers, it remains uncertain whether surface soil inputs provide a significant C source at 7 m depth.

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